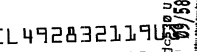
Page 1 of 2

#### CERTIFICATE OF EXPRESS MAILING

I hereby certify that this correspondence is being deposited with the United States Postal Service, Express Mail # 2832119US, postage prepaid, in an envelope addressed to Box Patent Application, Assistant Commissioner for Pater ington, D.C. 20231-9999 on June 1, 2000.

June 1, 2000





#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

# REQUEST FOR FILING APPLICATION

Under Rule 53(a), (b) & (f)

(No Filing Fee or Oath/Declaration) (Do NOT use for Provisional or PCT Applications) Use for Design or Utility Applications

**PATENT** APPLICATION

# **RULE 53(f) NO DECLARATION**

Assistant Commissioner of Patents	Atty. Dkt.	PMS-260755	
and Trademarks	·	M#	Client Ref
Washington, DC 20231 Box Patent Application			
• •	Date:	June 1, 20	000
Sir.		·	
This is a Request for filing a new Patent Application	Decian MI	Itility) antitlad:	
This is a Request for hing a new raterit Application	] Design 🖂 C	Itility) entitled:	
2 (Complete) Title: ANTIFUNGAL AND AN	TIMVCOBACTI	PIAL BASILISK	AMIDES
	TIMITOODAGTI	INIAL DAGILISH	CAMIDLS
without a filing fee or Oath/De	claration but for	which is enclose	d the following:
3 Abstract 2 page(s).			
Pages of Specification (only spec. and clai	ms): 5. 🗀 Spe	ecification in non-	English language
6 15 Numbered claim(s); and	,,		
7 Drawings: sheet(s)	et informal; 8.	formal of size	e:
a DOMEOTIC/INTERNATIONAL			
DOMESTIC/INTERNATIONAL priority is claimed u     following provisional and/or PCT int	inder 35 USC 1	9(e)/120/365(c)	based on the
following provisional, nonprovisional and/or PCT int	ernational applic	ation(s):	
9. DOMESTIC/INTERNATIONAL priority is claimed to following provisional, nonprovisional and/or PCT int Application No. Filing Date (1) 60/137,166 06/01/99	ernational applicat  Applicat (2)	ation(s):	based on the Filing Date
following provisional, nonprovisional and/or PCT int  Application No. Filing Date	ernational applicat Applicat (2)	cation(s):	
following provisional, nonprovisional and/or PCT int  Application No. Filing Date  (1) 60/137,166 06/01/99	ernational applicat Applicat (2)	eation(s): ion No.	
following provisional, nonprovisional and/or PCT int  Application No. Filling Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)	ernational applicat Applicat (2) -(d)/365(b) base	eation(s): ion No.	Filing Date
following provisional, nonprovisional and/or PCT int  Application No. Filling Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1) (No.) Certified copy (copies): attached	ernational applicat (2) -(d)/365(b) base Applicat (2) d; previo	eation(s): ion No.	Filing Date
following provisional, nonprovisional and/or PCT int  Application No. Filing Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1)  11. (No.) Certified copy (copies): attached in U.S. Application No. /	ernational applicat (2) -(d)/365(b) base Applicat (2) d; previo	eation(s): ion No.	Filing Date
following provisional, nonprovisional and/or PCT int  Application No. Filling Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1)  11. (No.) Certified copy (copies): attached in U.S. Application No. /  12. This is a reissue of Patent No.	ernational applicat (2)(d)/365(b) base Applicat (2) d;	eation(s): ion No. ed on filing in ion No. usly filed (date)	Filing Date Filing Date
following provisional, nonprovisional and/or PCT int  Application No. Filing Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1)  11. (No.) Certified copy (copies): attached in U.S. Application No. /  12. This is a reissue of Patent No.  13. See top first page re prior Provisional, National there and do not complete corresponding item	Applicat (2) -(d)/365(b) base Applicat (2) d;	eation(s):  ion No.  ed on filing in  ion No.  usly filed (date)  application(s) (X i	Filing Date Filing Date box only if info is
following provisional, nonprovisional and/or PCT int  Application No. Filling Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1)  11. (No.) Certified copy (copies): attached in U.S. Application No. /  12. This is a reissue of Patent No.  13. See top first page re prior Provisional, National	Applicat  (2)  -(d)/365(b) base  Applicat  (2)  d;	eation(s):  ion No.  ed on filing in  ion No.  usly filed (date)  application(s) (X in this is a  Co	Filing Date  Filing Date  box only if info is  ontinuation-in-Part
following provisional, nonprovisional and/or PCT int  Application No. Filing Date  (1) 60/137,166 06/01/99  10. FOREIGN priority is claimed under 35 USC 119(a)  Application No. Filing Date  (1)  11. (No.) Certified copy (copies): attached in U.S. Application No. /  12. This is a reissue of Patent No.  13. See top first page re prior Provisional, National there and do not complete corresponding item  14. Amend the specification by inserting before	Applicat  (2)  -(d)/365(b) base  Applicat  (2)  d;	eation(s):  ion No.  ed on filing in  ion No.  usly filed (date)  application(s) (X in this is a  Co	Filing Date  Filing Date  box only if info is  ontinuation-in-Part

15. Amend the s	<b>specification</b> by i enefit of U.S. Pro	nserting before visional Applica	e the first line:This ation No. 60/_137,166	application 6 , filed_06/01/99
16. Extension to date	: Concurrent	ly filed 🛛 r	not needed  pre	eviously filed
17. Prior application	is assigned to			
by Assignment recorded			Reel	Frame
18. Attached: Pre	liminary Amendm	ent		
19. This application is made inventor(s)	by the following na	amed	(Double check inst	ructions for accuracy.):
(1) Inventor Michael	i	T.	Kelly	
	First	Middle Initial		Family Name
Residence Surrey, British	n Columbia	Canada		Canadian
	City	State	e/Foreign Country	Country of Citizenship
Post Office Address	1825 - 133A Str	reet, Surrey, B	ritish Columbia, Cana	ada
(include Zip Code)	V4A 7M4			
	<del></del>	<del></del>		
(2) Inventor Raymond		J.	Andersen	
	First	Middle Initial		Family Name
Residence Vancouver, B	ritish Columbia	Canada	<del></del>	Canadian
	City		e/Foreign Country	Country of Citizenship
Post Office Address			couver, British Colun	
(include Zip Code)	V6S 1Z6	1	oddror, British Goldin	insia, Gariada
Bendit	1 1 3 1 2 3			
(3) Inventor Todd		Α.	Barsby	
	First	Middle Initial		Family Name
E a face of the face	ritish Columbia	Canada		Canadian
e a company de la company	City	<del></del>	e/Foreign Country	· · · · · · · · · · · · · · · · · · ·
Post Office Address			couver, British Colum	Country of Citizenship
(include Zip Code)	V6T 2W5	T	Boaver, British Colum	iola, Carlada
Supplied State of the State of	V01 2VV3	_		
grander G				
20. NOTE: FOR ADDITION	ONAL INVENTOR	RS check box		
and attach sheet wi				
		Pillsbury Madison		
		ntellectual Prope		
50 Fremont Street Fifth Floor	By: Atty:	Georgina M. McPau	1	Reg. No. <u>42,873</u>
San Francisco, CA 94105-2230 Tel: (415) 983-1000	Sig:	GM De	and a	Fax: (415) 983-1200
Atty/Sec: GMM/gfp				Tel: (415) 983-1718
	NOTE: File in duplicat	te with 2 post card r	receipts (PAT-103) & attach	ments

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Non Provisional Patent Application of	Group Art Unit: Unassigned
Michael T. Kelly, et al.	Examiner: Unassigned
Filed: June 1, 2000	PRELIMINARY AMENDMENT
For: ANTIFUNGAL AND ANTIMYCOBACTERIAL BASILISKAMIDES	) ) ) )

#### CERTIFICATE OF EXPRESS MAILING

I hereby certify that this correspondence is being deposited with the United States Postal Service, Express Mail # EL492832119US, postage prepaid, in an envelope addressed to Box Patent Application, Assistant Commissioner for Patents, Washington, D.C. 20231-9999 on June 1, 2000.

June 1, 2000

Guy Powers

Assistant Commissioner for Patents and Trademarks Washington, D.C. 20231 Box Patent Application

Sir:

Entry of the following preliminary claim amendments is respectfully requested in the subject application. No new matter has been added.

#### In the Claims:

Please cancel claims 15 and 16.

Please amend claims 3, 7-11 and 14 as follows:

3. (Amended) The compound or physiologically acceptable salt thereof of claim 1 ex 2 wherein Z' is a linear or branched, saturated or unsaturated one to eight carbon carbonyl optionally substituted with a substituent selected from the group consisting of: NH<sub>2</sub>, NHR, NR<sub>2</sub>, OH, OR, SH, SR, H and CF<sub>3</sub>, wherein R is as defined.

7. (Amended) The compound or physiological salt thereof of any one of claims 4-6 claim 4, wherein  $R_1$  and  $R_2$  are independently H or  $CH_3$ .

8. (Amended) The compound or physiological salt thereof of any one of claims 4-6 claim 4, wherein  $R_3$  is (a)

9. (Amended) The compound or physiological salt thereof of any one of claims 4-6 claim 4, wherein X is NH<sub>2</sub>.

10. (Amended) The compound or physiological salt thereof of any one of claims 4-6 claim 4, wherein  $R_3$  at  $C_7$  is (a) and  $R_3$  at  $C_9$  is OH.

11. (Amended) The compound or physiological salt thereof of any one of claims 4-6 claim 4, wherein  $R_3$  at  $C_7$  is OH and  $R_3$  at  $C_9$  is (a).

14. (Amended) A pharmaceutical composition comprising a compound or physiological salt thereof of any one of claims 1-13 claim 1, and a pharmaceutically acceptable carrier.

Please add the following new claim:

--17. A pharmaceutical composition comprising a compound or physiological salt thereof of claim 4, and a pharmaceutically acceptable carrier.

Respectfully submitted,

Dated: June 1, 2000

Georgina M. McPau

Reg. No. 42,873

PILLSBURY MADISON & SUTRO LLP 50 Fremont Street

San Francisco, CA 94105 Telephone: 415/983-1718

Fax: 415/983-1200

# 5 ANTIFUNGAL AND ANTIMYCOBACTERIAL BASILISKAMIDES

#### Field of the Invention

This invention relates to polyketide amides having antibiotic activity.

10

15

20

25

#### **Background of the Invention**

There is an urgent need for new antibiotics to treat pathogens that have developed resistance to antibiotics currently in use. Further, compounds that have antimycobacterial activity are rare. Compounds produced by marine microorganisms are being screened for antibiotic activity.

Japanese patent application 06-27802 published September 12, 1995 under No. 07238018 and entitled "Antimycotic Antibiotic Substance and its Production" discloses an antifungal compound YL-03709B-A obtained by fermentation of *Bacillus sp.* YL-03709B (FERM P-14126). The *Bacillus* was isolated from soils near Okinawa, Japan. The compound was reported as having antifungal activity against several organisms but low activity against *Candida albicans*, *Candida parapellosis*; *Saccharomyces cerevisiae*; *Saccharomyces sake*; and *Aspergillus niger* on Sabouraud/dextrose Agar medium.

An unidentified *Bacillus sp.* (MK-PNG-276A) was isolated from the tissues of a tubeworm collected in the tropical waters off Papau, New Guinea. Extracts from laboratory cultures of the latter organism exhibited broad spectrum antibiotic activity against a panel of antibiotic-resistant pathogens. Initial bioassay guided fractionation of crude extracts resulted in the isolation of the loloatins, a family of novel cyclic decapeptides (see PCT/CA97/00529). More recently, a class of novel polyketide amides were isolated from MK-PNG-276A cultures, which are termed basiliskamides herein. The basiliskamides have antibiotic activity.

## **Summary of the Invention**

This invention provides a compound or a physiologically acceptable salt thereof, wherein the compound has the formula:

10

$$Z^{1}$$
 $R_{3}$ 
 $R_{4}$ 
 $R_{2}$ 
 $R_{4}$ 

15

wherein:

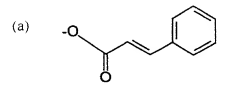
 $R_1$  and  $R_2$  are the same or different and are independently H or R;

20

25

R is a structural fragment having a saturated or unsaturated linear, branched, or cyclic, skeleton containing one to ten carbon atoms in which the carbon atoms may be optionally substituted with a substituent selected from the group consisting of: -OH; =O; -OR $_5$ ; -O $_2$ CR $_5$ , -SH; -SR $_5$ ; -SOCR $_5$ ; -NH $_2$ ; -NHR $_5$ ; -NH(R $_5$ ) $_2$ ; -NHCOR $_5$ ; NRCOR $_5$ ; -I: -Br; -Cl; -F; -CN; -CO $_2$ H; -CO $_2$ R $_5$ ; -CHO; -COR $_5$ ; -CONH $_2$ ; -CONHR $_5$ ; -CON(R $_5$ ) $_2$ ; -COSH: -COSR $_5$ ; -NO $_2$ ; -SO $_3$ H; -SOR $_5$ ; and -SO $_2$ R $_5$ , wherein R $_5$  is a linear, branched or cyclic, one to ten carbon saturated or unsaturated alkyl group;

 $R_3$  and  $R_4$  are different and are independently selected from the groups consisting of OH,



20

and

$$(b)$$
  $-O-Z-Ar$ 

wherein,

10 Z<sup>1</sup> and Z are linear or branched, saturated or unsaturated, one to ten carbon fragments optionally substituted with Y;

Ar is a monocyclic, bicyclic or tricyclic, fully or partially aromatic system containing five or six membered carbocyclic or, oxygen, nitrogen or sulphur containing heterocyclic rings, optionally substituted with R or Y;

Y is selected from the group consisting of: H; =O, -OH: -OR; -O<sub>2</sub>CR; -SH; -SR; -SOCR; -NH<sub>2</sub>; -NHR; -NH(R)<sub>2</sub>; -NHCOR; NRCOR; -I; -Br; -Cl; -F; -CN- -CO<sub>2</sub>H: -CO<sub>2</sub>R; -CHO; -COR; -CONH<sub>2</sub>; -CONHR; -CON(R)<sub>2</sub>; -COSH; -COSR; -NO<sub>2</sub>; -SO<sub>3</sub>H: -SOR; -SO<sub>2</sub>R; and, -O- (epoxide);

W is H or R;

with the provisos that when W is H,  $R_2$  is not H; when  $R_2$  is  $CH_3$ , W is not n-propyl: 25 and, one of  $R_3$  and  $R_4$  is (a) or (b) and another of  $R_3$  and  $R_4$  is OH.

This invention also provides a compound or a physiologically acceptable salt thereof, wherein the compound has the formula:

30 
$$C_1$$
  $C_2$   $C_3$   $C_4$   $C_5$   $C_6$   $C_7$   $C_8$   $C_{10}$   $C_{11}$   $C_{12}$   $C_{12}$ 

wherein:

a single, double or triple bond exists between one or more of: C-2 and C-3: C-3 and C-4; C-4 and C-5; and, C-5 and C-6;

X is NH<sub>2</sub>, NHR, NR<sub>2</sub>. OH, OR, SH, SR, H, or CF<sub>3</sub>:

R is a structural fragment having a saturated or unsaturated linear, branched, or cyclic. skeleton containing one to ten carbon atoms in which the carbon atoms may be optionally substituted with a substituent selected from the group consisting of: -OH; =O; -OR5; -O2CR5, -SH; -SR5; -SOCR5; -NH2; -NHR5; -NH(R5)2; -NHCOR5; NRCOR5; -I; -Br; -Cl; -F; -CN; -CO2H; -CO2R5; -CHO; -COR5; -CONH2; -CONHR5; -CON(R5)2; -COSH: -COSR5; -NO2; -SO3H; -SOR5: and -SO2R5, wherein R5 is a linear, branched or cyclic, one to ten carbon saturated or unsaturated alkyl group;

 $R_1$  and  $R_2$  are the same or different and are independently H or R;

25 R<sub>3</sub> and R<sub>4</sub> are different and are selected from the group consisting of: OH.

and

1 ...

15

20

25

30

35

wherein, Z is a linear or branched, saturated or unsaturated, one to ten carbon fragment optionally substituted with Y;

Ar is a monocyclic, bicyclic or tricyclic, fully or partially aromatic system containing five or six membered carbocyclic or, oxygen, nitrogen or sulphur containing heterocyclic rings, optionally substituted with R or Y;

Y is selected from the group consisting of: H; =O, -OH; -OR; -O<sub>2</sub>CR; -SH; -SR; -SOCR: -NH<sub>2</sub>; -NHR; -NH(R)<sub>2</sub>; -NHCOR; NRCOR; -I; -Br; -Cl; -F; -CN- -CO<sub>2</sub>H; -CO<sub>2</sub>R: -CHO; -COR: -CONH<sub>2</sub>; -CONHR; -CON(R)<sub>2</sub>; -COSH; -COSR; -NO<sub>2</sub>; -SO<sub>3</sub>H; -SOR: -SO<sub>2</sub>R: and. -O- (epoxide);

with the proviso that one of  $R_3$  and  $R_4$  is (a) or (b), and another of  $R_3$  and  $R_4$  is OH.

If a compound of this invention is naturally occurring (such as basiliskamide A or B as described herein) such a compound may be obtained from a natural source or may be sythnesized as described herein. In cases where such a naturally occurring compound is obtained from a natural source, the compound of this invention is characterized as being purified or partially purified. Thus, any compound of this invention that is naturally occurring will be substantially free of cellular contaminants. Cellular contaminants are defined as any component of a living cell (eg. proteins, nucleic acids, cell wall fragments, etc.) or a naturally occurring compound that is not a compound of this invention. The term "substantially free of cellular contaminants" means that a compound or a mixture of compounds of this invention, whether or not present in a pharmaceutical composition, will be present at a ratio of at least 3:1 (w/w) of the total amount of a compound or compounds of this invention to total amount of cellular contaminants present.

This invention also provides pharmaceutical compositions comprising a compound of this invention and a pharmaceutically acceptable carrier selected for the particular indication and mode of treatment in which the compound is to be used.

Compounds of this invention that are capable of forming salts may be in the form of a physiologically acceptable salt. Such a salt is any salt that is acceptable for use in pharmaceutical formulations. For example, where a compound of this invention has a carboxyl or sulphonic acid moiety, the counterion may be Na, K, Mg or Zn. Where the compound comprises a basic moiety such as an amine, the salt may be hydrochloride.

Pharmaceutical compositions of this invention will contain a compound or physiologically acceptable salt thereof in admixture with any carrier, excipient, dilutant, filler, thickener, etc., or in combination with any drug delivery moiety or device selected as suitable for a particular indication and mode of treatment desired. For example, pharmaceutical compositions of this invention may be formulated for injection (intravenous or otherwise), topical application, oral dosage (eg. tablets, capsules, powders), eye drops, aerosol delivery or cosmetic/cleansing formulations such as shampoos or skin cleansers. Selection of pharmaceutically acceptable carriers for compounds of this invention are within the knowledge of those of skill in the art. An example of a formulation for topical use is a creme-based formulation or carrier in which a compound of this invention or a pharmaceutically acceptable thereof is dissolved or emulsified.

This invention also provides a method for treatment of a patient (animal or human) afflicted with a fungal or mycobacterial infection comprising the administration to said patient of a therapeutically effective amount or a compound of pharmaceutical composition of this invention. This invention provides the use of a compound or pharmaceutical composition of this invention as an antifungal agent or as an antimycobacterial agent. The dose range of a compound of this invention will be selected in accordance with a particular indication or mode of treatment. Generally, the dose range will be between about 50 and about 500 mg/day for oral and intravenous applications and between about 0.5 and about 5 gram for topical applications.

Marine Bacterium MK-PNG-276A: The marine bacterium MK-PNG-276A was isolated during a collecting expedition off of Loloata Island, Papua New Guinea. MIDI analysis of cellular fatty acids indicated that MK-PNG-276A was an unknown species possibly within the genus *Bacillus*. MK-PNG-276A was deposited July 2. 1996 at the American Type Culture Collection (ATCC) under No. 55797.

Isolation of the Basiliskamides: The marine bacterium MK-PNG-276A was grown in moderate scale culture as confluent lawns for 5 days at 16 °C on trays of solid trypticase soy agar supplemented with NaCl to a final concentration of 1%. The cultures were harvested by gently scraping the cells from the agar surface. Bacterial cells (21.5 g dry weight) were immersed in and subsequently extracted with MeOH (3 X 250 mL) over a period of six days. Crude MeOH extracts showed broad spectrum antimicrobial activity against a variety of human pathogens, including methicillin resistant Staphylococcus aureus, Eschericia coli, Candida albicans and Mycobacterium tuberculosis.

The combined MeOH extracts were concentrated in vacuo and then partitioned between EtOAc (3 x 100 mL) and H<sub>2</sub>O/MeOH (10:1 200 mL). The EtOAc extract was dried over anhydrous Na<sub>2</sub>SO<sub>4</sub>, filtered and reduced to dryness in vacuo to give 6.5 g of a gum. The gum was fractionated by Sephadex LH-20 chromatography (eluent MeOH) to give 226 mg of a fraction containing strongly UV absorbing compounds. This fraction was subsequently subjected to step gradient reversed-phase chromatography (eluent: 1:1 MeOH/H<sub>2</sub>O to 100% MeOH) on a 10g Waters Sep-Pak. A strongly UV absorbing fraction (82 mg) was further separated into crude basiliskamide A and crude basiliskamide B (28 mg total) by a normal-phase silica gel flash chromatography (4:1 EtOAC/CH<sub>2</sub>Cl<sub>2</sub>). Final purification was accomplished by reversed-phase HPLC (7:3 MeOH/H<sub>2</sub>O), yielding pure basiliskamides A (1, 14 mg) and B (2, 9 mg) as clear solids.

Structure Elucidation of Basiliskamides A (1) and B (2): Basiliskamide A (1) was isolated as a clear solid that gave a  $[M + H]^+$  ion at m/z 386.23358 in the high resolution fast atom bombardment mass spectrum appropriate for a molecular formula

of C<sub>23</sub>H<sub>31</sub>NO<sub>4</sub>. The <sup>13</sup>C NMR spectrum (Table 1) of basiliskamide A (1) showed only 21 well resolved resonances, indicating that there was an element of symmetry in the molecule. Resonances in the <sup>1</sup>H NMR spectrum of basiliskamide A were all well dispersed, which facilitated identification of the two major substructures.

A broad three proton  $^1$ H NMR resonance at  $\delta$  7.40-7.41. that showed HMQC correlations to carbon resonances at  $\delta$  128.9 and 130.4, along with a broad two proton  $^1$ H NMR resonance at  $\delta$  7.71, that showed HMQC correlations to a carbon resonance at  $\delta$  128.4, were all assigned to a monosubstituted phenyl ring. The phenyl ring accounted for the element of symmetry required by the  $^{13}$ C NMR data. A one proton doublet at  $\delta$  7.65 in the  $^1$ H NMR spectrum showed COSY correlations to the phenyl multiplet at  $\delta$  7.71 and to another one proton doublet at  $\delta$  6.61. The two doublets were assigned to a vinyl group that was the only substituent on the phenyl ring. HMBC correlations observed between the vinyl doublet resonance at  $\delta$  7.65 and the phenyl ring. HMBC correlations observed between both of the vinyl group to the phenyl ring. HMBC correlations observed between both of the vinyl proton resonances at  $\delta$  7.65 and 6.61 and a carbon resonance at  $\delta$  166.0, showed that the phenyl and vinyl fragments were part of a cinnamoyl residue. The vinyl protons had a vicinal scalar coupling of 16 Hz demonstrating the cinnamoyl residue had the E configuration.

Analysis of COSY, HMQC, and HMBC data collected for basiliskamide A (1) routinely identified the linear carbon chain extending from C-2 to C-12, including the positions of the  $\Delta^{2.3}$  and  $\Delta^{4.5}$  olefins, the methyl branches at C-8 and C-10, and the presence of -OR substituents at C-7 and C-9. HMBC correlations observed between both the H-2 and H-3 resonances at  $\delta$  5.55 and 6.31, respectively, and a carbon resonance at  $\delta$  167.5, showed that C-2 was attached to a carbonyl carbon. Only one nitrogen and two hydrogen atoms remained unaccounted for by the cinnamoyl and linear C-1 to C-12 chain fragments, suggesting that the C-1 carbonyl was a primary amide. A pair of broad one proton resonances at  $\delta$  6.82 and 7.32, that showed COSY correlations to each other but did not show HMQC correlations to carbon resonances. were assigned to the primary amide NH protons. The NH resonance at  $\delta$  6.82 showed an HMBC correlation to the C-2 resonance at  $\delta$  119.3, confirming the presence of the primary amide at the terminus of the linear C-1 to C-12 carbon chain. A COSY

20

25

30

35

5 correlation observed between an OH proton resonance at δ 4.57 and the H-7 resonance at δ 3.55 showed that there was an alcohol functionality at H-7 and, therefore, the cinnamoyl fragment had to be attached to the linear carbon chain via an ester linkage at C-9. An HMBC correlation observed between the H-9 methine resonance at δ 4.92 and the cinnamoyl carbonyl resonance at δ 166.0 confirmed the presence of the C-9 ester linkage.

H-2 and H-3 had a vicinal scalar coupling constant of 11 Hz typical of Z olefins, while H-4 and H-5 showed a 15 Hz vicinal coupling typical of E olefins. Difference nOe experiments confirmed the assigned olefinic configurations. Irradiation of the H-3 resonance at  $\delta$  6.31 induced an nOe in the H-2 resonance at  $\delta$  5.55 in agreement with the Z configuration for the  $\Delta^{2.3}$  olefin. Similarly, irradiation of the H-5 resonance at  $\delta$  5.91 induced a strong nOe in the H-3 resonance at  $\delta$  6.31 supporting the E configuration for the  $\Delta^{4.5}$  olefin.

The relative stereochemistry at C-7 and C-9 was determined by converting basiliskamide A (1) to the acetonide derivative 7. Analysis of the HMQC data for 7, showed that the acetonide methyl carbon resonances had chemical shifts of 19.8 and 30.4 ppm, typical of acetonides formed from syn-1,3-diols. Further analysis of the <sup>1</sup>H NMR data for the acetonide 7 showed that the dioxane ring existed in a chair conformation with the C-6 and C-10 carbons equatorial. A vicinal coupling constant of 10 Hz was observed between H-9 and H-8 indicating that H-8 was axial and. therefore, the C-14 methyl had to be equatorial, establishing the relative stereochemistries at C-7, C-8, and C-9 as shown in 7. Standard Mosher ester methodology was used to show that C-7 in basiliskamide A (1) had the S configuration. The configuration of C-10 in 1 was not determined. However, basiliskamide A (1) is a homolog of YM47522 (5) and the absolute configuration at C-10 in 5 has been determined by synthesis to be R. Since the other chiral centers in 1 and 5 have identical configurations, and the and the <sup>1</sup>H and <sup>13</sup>C NMR data for 1 and 5 are nearly identical for the C-7, C-8, C-9 and C-10 centers, it is indicated that 1 also has the R configuration at C-10.

Basiliskamide B (2) was also isolated as a clear solid that gave a  $[M + H]^+$  ion at m/z 386.23358 in the high resolution fast atom bombardment mass spectrum appropriate for a molecular formula of  $C_{23}H_{31}NO_4$ , identical to the formula of

basiliskamide A (1). Analysis of the 1D and 2D NMR data obtained for basiliskamide B (2) showed that it was simply an isomer of basiliskamide A, in which the cinnamoyl ester was at C-7 instead of C-9. Basiliskamide B (2) and basiliskamide A (1) were both converted to the same diol 6 by DIBAL reduction, demonstrating that both molecules had identical absolute configurations.

Basiliskamide A (1): isolated as a clear solid; <sup>1</sup>H NMR, see Table 1; <sup>13</sup>C NMR, see Table 1; IR (film)  $\upsilon_{max}$ : 3348, 3205, 2966, 2934, 1705, 1697, 1635, 1595, 1450 cm<sup>-1</sup>; UV (MeOH)  $\lambda_{max}$ : 262 nm ( $\varepsilon$  41 000); [ $\alpha$ ]<sup>25</sup>D (MeOH) = -78; pos itive-ion HRFABMS [M+ H]<sup>+</sup> m/z 386.23358 (C<sub>23</sub>H<sub>32</sub>NO<sub>4</sub>, calcd 386.23313).

Basiliskamide B (2): isolated as a clear solid; <sup>1</sup>H NMR, see Table 1; <sup>13</sup>C NMR, see Table 1; IR (film)  $v_{max}$ : 3348, 3205, 2962, 2926, 1702, 1664, 1637, 1595, 1450 cm<sup>-1</sup>; UV (MeOH)  $\lambda_{max}$ : 262 nm ( $\epsilon$  43 000);  $[\alpha]^{25}_D$  (MeOH) = -12; HREIMS  $[M]^+$  m/z 385,22531 (C<sub>23</sub>H<sub>31</sub>NO<sub>4</sub>, calcd 385,22531).

Reduction of Basiliskamides. To basiliskamides B (2, 4.7 mg) in 1 mL

THF under Ar (g), at -78, 4 equivalents of diisobutylaluminum hydride (DIBAL-H) were added. The reaction was stirred overnight then diluted with EtOAc (3 mL) and 20 quenched by the addition of 2 mL NH4Cl (aq), stirring until the reaction mixture turned cloudy (10 min). The mixture was extracted thrice with EtOAc, and the combined organics were reduced to dryness in vacuo. Preparative normal-phase TLC (100 % EtOAc) followed by reversed-phas HPLC (70/30 MeOH/H<sub>2</sub>O, 280 nm) gave 2 mg of 6. <sup>1</sup>H NMR (500 MHz, DMSO-d6)  $\delta$  7.44 (1H, dd, J = 15 Hz, 11Hz), 7.35 25 (1H. br s, NH), 6.86 (1H, br s, NH), 6.36 (1H, dd, J = 11 Hz), 6.02 (1H, dt, J = 15 Hz, 7 Hz), 5.57 (1H, d, J = 11 Hz), 4.48 (1H, d, J = 5.4 Hz, OH). 4.69 (1H, d, J = 4.4 Hz. OH), 3.80 (1H, m), 3.23 (1H, m), 2.26 (1H, m), 2.07 (1H, m), 1.61 (1H, m), 1.36 (1H, m). 1.35 (1H, m), 1.18 (1H, m), 0.84 (3H, t. J = 7 Hz), 0.72 (3H, d. J = 7 Hz), 0.66 (3H, d. J = 7 Hz); positive-ion HRFABMS [M+H]<sup>+</sup> m/z 256.19211 (C14H26NO3. 30 calcd 256.19127).

Formation of Acetonide (7). To 1.5 mg of 6 in 0.5 mL 2,2-dimethoxypropane, pyridinium p-toluenesulfonate (5 wt% diolbasiliskamide) was added. The reaction mixture was stirred under Ar (g) and heated at 60 C for 1 h.

25

The reaction mixture was filtered through silica (rinsed with EtoAc) and the solvents 5 removed in vacuo. Reversed-phase HPLC (80/20 MeOH/H2O) yielded 1 mg of 7. <sup>1</sup>H NMR (400 MHz, DMSO-d6)  $\delta$  7.46 (1H, dd, J = 15 Hz, 11 Hz, H-4), 7.35 (1H, br s. NH). 6.85 (1H, br s, NH), 6.37 (1H, dd, J = 11 Hz, 11 Hz, H-3), 5.94 (1H, dt, J = 15Hz, 7 Hz, H-5), 5.58 (1H, d, J = 11 Hz, H-2), 3.57 (1H, m, H-7), 3.48 (1H, dd, J = 10Hz. 2 Hz, H-9), 2.44 (1H, m, H-6), 2.19 (1H, m, H-6'), 1.54 (1H, m, H-10), 1.36 (3H. 10 s, Me-17), 1.33 (1H, m, H-8), 1.30 (1H, m, H-11), 1.25 (1H, m, H-11'), 1.23 (3H, s. Me-16), 0.83 (3H, t, J = 7 Hz, Me-12), 0.75 (3H, d, J = 7 Hz, Me-13), 0.71 (3H, d, J = 7 Hz, Me-14), 0.71 (3H, d, J = 7 Hz, Me-15), 0.71 (3H, d, J = 7 Hz, 7 Hz. Me-14); <sup>13</sup>C NMR (DMSO-d6, 100 MHz) δ 167.8 (C-1), 140.6 (C-3), 138.5 (C-5). 128.9 (C-4), 120.2 (C-2), 97.6 (C-15), 74.9 (C-9), 74.0 (C-7), 36.5 (C-6), 35.1 (C-8). 34.6 (C-10), 30.4 (C-16), 26.7 (C-11), 19.8 (C-17), 12.7 (C-13), 12.1 (C-12). 15 11.6 (C-14); positive-ion HRFABMS [M+ H]+ m/z 296.22198, C17H30NO3, calcd 296.2257.

Reaction of 1 with (*R*)-MTPA Acid. To a solution of 1 (1.5 mg) in 0.5 mL dry CH<sub>2</sub>Cl<sub>2</sub> were added DMAP (1 mg), a drop of triethylamine and (*R*)-MTPA acid (4 mg) and the solution stirred for 16 h. Removal of solvent in vacuo, followed by preparative reversed-phase TLC (100% MeOH), then reversed-phase HPLC (MeOH/H2O 4:1) gave the (*R*)-MTPA ester 1a (0.8 mg). <sup>1</sup>H NMR (500 MHz, DMSO-*d*6)  $\delta$  7.72 (3H, br envelope), 7.43 (9H, br envelope), 7.35 (1H, br s), 6.84 (1H, br s), 6.70 (1H, d, J = 16 Hz), 6.20 (1H, dd, J = 11 Hz, 11 Hz), 5.58 (2H, m), 5.17 (1H, m), 4.98 (1H, m), 3.43 (3H, s), 2.60 (1H, m), 2.26 (2H, br m), 1.72 (1H, m), 1.29 (2H, br m), 1.17 (1H, m), 0.95 (3H, d, J = 7 Hz), 0.93 (3H, d, J = 7 Hz), 0.88 (3H, t, J = 7 Hz); positive-ion HRFABMS [M+H]+ m/z 602.27148, C<sub>33</sub>H<sub>39</sub>NO<sub>6</sub>F<sub>3</sub>. calcd 602.272950.

Reaction of 1 with (S)-MTPA Acid. To a solution of 1 (1.5 mg) in 0.5 mL dry CH<sub>2</sub>Cl<sub>2</sub> were added DMAP (1 mg), a drop of triethylamine and (S)-MTPA acid (4 mg) and the solution stirred for 16 h. The reaction was quenched and purified as above, yielding the (S)-MTPA ester 1b (0.4 mg). <sup>1</sup>H NMR (500 MHz, DMSO-d6)  $\delta$  7.72 (3H, br envelope), 7.54 (1H, m), 7.43 (8H, br envelope), 7.38 (1H, br s), 6.91 (1H, br s), 6.72 (1H, d, J = 16 Hz), 6.36 (1H, dd, J = 11 Hz, 11 Hz), 5.78 (1H, m).

5 5.64 (1H, d, J = 11 Hz), 5.13 (1H, m), 4.94 (1H, m), 3.42 (3H, s), 2.63 (1H, br m), 2.35 (1H, m), 2.17 (1H, m), 1.65 (1H, m), 1.27 (2H, br m), 1.15 (1H, m), 0.89 (3H, d, J = 7 Hz), 0.87 (3H, t, J = 7 Hz), 0.70 (3H, d, J = 7 Hz); positive-ion HRFABMS [M+H]<sup>+</sup> m/z 602.27352, C33H39NO6F3, calcd 602.272950.

**Table 1.** <sup>13</sup>C (100 MHz) and <sup>1</sup>H (500 MHz) NMR Spectal Data for Basiliskamides A and B in DMSO-d<sub>6</sub>

Basiliskamide A				Basiliskamide B				
(1)				(2	)			
Atom	δ <sup>13</sup> C	δ <sup>1</sup> H <i>J</i> (Hz))	(intgrn,	m,	δ <sup>13</sup> C	δ <sup>1</sup> H <i>J</i> (Hz))	(intgrn,	m,
1	167.5				167.4			
2	119.3	5.55 (1)	H, d, 11)		119.9	5.57 (1	H, d, 11)	
3	140.5		H, dd, 11, 1	1)	140.0		H, dd, 11,	11)
4	128.2	7.40 (1			128.8		H, dd, 15,	•
5	140.5	,	H, dt, 15, 7	)	138.0		H, dt, 15, 7	
6	34.7	2.28 (1			31.8	2.53 (1		
6'		1.99 (1	H, m)			2.36 (1	-	
7	69.6	3.49 (1			73.0	-	H, dt, 10.5	, 3)
8	40.7	2.03 (1	H, m)		39.4	1.92 (1	H, m)	
9	76.3	4.92 (1	H, dd, 9.5,	2)	74.0		H, m)	
10	35.5	1.67 (1	H, m)		36.3	1.40 (1	H, m)	
11	26.4	1.25 (1	H, m)		26.5	1.38 (1	H, m)	
11'		1.11 (1	H, m)			1.21 (	lH, m)	
12	10.1	0.87 (3	SH, t, 7.5)		11.8	0.85 (	3H, t, 7)	
13	11.6	0.84 (3	3H, d. 7)		10.7	0.83 (	3H, d, 7)	
14	12.8	0.90 (3	SH, d, 7)		12.1	0.74 (	3H, d, 7)	
15	166.0				165.5			
16	118.0	6.61 (	1H, d, 16)		118.5	6.59 (	1H, d, 16)	
17	144.6	7.65 (	1H, d, 16)		144.1	7.60 (	1H, d, 16)	
18	134.0				134.0			
19	128.4	7.71 (	2H, m)		128.2	7.70 (	2H, m)	
20	128.9	7.41 (	2H. m)		129.0	7.40 (	2H, m)	
21	130.4	7.40 (	1H. m)		130.2	7.40 (	1H, m)	
$NH_2$		7.31,	6.83 (2H, s)	<b>)</b>		7.34,	6.86 (2H. s	)
ОН		4.57 (	1H, d, 5)			4.48 (	(1H, m)	

# Basiliskamide A

# Basiliskamide B

$$H_2N$$
 $7$ 
 $\overline{0}$ 
 $\overline{0}$ 

Preparation of Basiliskamides: Compounds of this invention may be prepared from a natural source by fermentation as described above, or by total synthesis, for example by modification of the total synthesis of YM47522 that was described in Ermokenko. M.S. Tetrahedron Letters, 1996, 37, 6711-12 (as exemplified in the scheme below for basiliskamide A).

- a)  $(CH_3CH_2)_2Mg/Et_2O$ ,  $\Delta$ , 0.5 h:
- a) NMO-Pr<sub>4</sub>NRuO<sub>4</sub> (0.02 eq), MS 4 Å/ MeCN, rt, 0.5 h;
- 15 b) NaBH<sub>4</sub>-CeCl<sub>3</sub>7H<sub>2</sub>O/MeOH, -20 C, 0.5 h;
  - c) HSCH<sub>2</sub>CH<sub>2</sub>SH, BF<sub>3</sub>.Et<sub>2</sub>O/CH<sub>2</sub>Cl<sub>2</sub>, rt, 2 h;
  - d)  $Ac_2O/Py$ , -10 C, 2h;
  - e) Me<sub>2</sub>CO-Me<sub>2</sub>C(OMe)<sub>2</sub>, TsOH (cat);
  - f) Ra-Ni/EtOH,  $\Delta$ , 0.5 h;

- 5 g) K<sub>2</sub>CO<sub>3</sub>/MeOH, rt, 0.5 h
  - h) TsCl/Py;
  - i) LiI-HMPA/PhMe,  $\Delta$ . 0.5 h;
  - j) Me<sub>2</sub>CuLi-LiCN, 2 eq (E)-Bu<sub>3</sub>SnCH=CHSnBu<sub>3</sub>/THF-Et<sub>2</sub>O, rt, 2 h, then 15. -78 C to rt, then NIS, rt;
- 10 k) (Z)-Bu<sub>3</sub>SnCH=CHCONH<sub>2</sub>, (MeCN)<sub>2</sub>PdCl<sub>2</sub> (0.05 eq)/DMF, rt, 24 h;
  - 1) AcOH-H<sub>2</sub>O (4:1), 60 C, 6 h;
  - m) 1.2 eq Et<sub>3</sub>SiCl/Py, 0 C, then PhCH=CHCOCl, DMAP/CH<sub>2</sub>Cl<sub>2</sub>, rt;
  - n) HF (aq)-MeCN, rt, 1 h.
- 15 Conversion of compound 10 to compound 11 is as described in Sviridov, A. F. et al. *Izv. Akad. Nauk SSR, Ser. Khim.* **1982**, 2572-2574.

Analogs can be prepared by modification of the total synthesis shown above or by semisynthesis from a product of the total synthesis or a naturally derived product. An example employing basiliskamide A (1) is shown in the scheme below.

$$\begin{array}{c|c} O & NH_2 \\ \hline O & O & NH_2 \\ \hline O & O & O & O \\ \hline O & O & O \\ \hline O & O & O & O$$

- a) DIBAL (4 eq), THF. -78 C, 15h
- b) 1.2 eq Et<sub>3</sub>SiCl/Py, 0 C
- c) alkylate or acylate alcohol (i.e for acylation ArCH=CHCOCl, DMAP/CH $_2$ Cl $_2$ , rt)
- d) HF (aq)-MeCN, rt, 1 h

10

25

Antifungal Activity of the Basiliskamides: The antifungal activity of basiliskamides A and B was compared to that of the known antifungal agent, amphotericin B. A standardized macrobroth dilution method as used which was developed and published by the National Committee for Clinical Laboratory Standards (Reference Method for Broth Dilution Antifungal Susceptibility Testing of Yeasts, Approved Standard 1996 M27-A, Vol. 15, No.10). An agar method based on the standardized broth method was also used. The results are presented in Tables 2-5 below.

Table 2. Antifungal Activity of Basiliskamide A and B
Tested by Macrobroth Dilution

## Minimal Inhibitory Concentration (µg/ml)

15	Basiliskamide	Candida albicans	Trichophyton rubrum	Aspergillus fumigatus
	Α	0.5	1.0	4.0
	В	≥32	4.0	≥32
20				

Table 3. Antifungal Activity of Basiliskamide A and B
Tested by Agar Dilution

# Minimal Inhibitory Concentration (µg/ml)

30	Basiliskamide	Candida albicans	Trichophyton rubrum	Aspergillus fumigatus
	A	1.0	0.5	2.5
	B	3.1	5.0	5.0

Table 4. Activity of Basiliskamide A Compared to Amphotericin B
Against 7 Clinical Isolates of Candida albicans as
Determined by Macrobroth Dilution

#### Minimal Inhibitory Concentration (µg/ml)

10	Isolate Number	Basiliskamide A	Amphotericin B
	8167	0.5	0.5
	8362	0.5	0.5
	8363	0.5	0.5
15	8364	0.5	0.5
	8365	0.5	0.5
	8366	0.5	0.5
	8367	0.5	0.5

20

Table 5. Comparative Activity of Basiliskamides A, B and YL-03709B-A as Determined by Agar Dilution

25

### Minimal Inhibitory Concentration (µg/ml)

2.0	Target Organism	Basiliskamide A	Basiliskamide B	Reported Value for YL-03709B-A
30	Candida albicans	1.0	3.1	25
	Aspergillus fumigatus	s 2.5	5.0	≥50

35

40

These findings demonstrate that the basiliskamides have superior antifungal activities and are active against the dermatophyte, *Trichophyton rubrum*, the yeast, *Candida albicans*, and the opportunistic fungus, *Aspergillus fumigatus*. Basiliskamide A activity against clinical isolates of the yeast, *Candida albicans* when tested by the broth dilution method is comparable to that of amphotericin B, a commonly used antifungal agent. The data presented in Table 5 demonstrates that Basiliskamide A is about 25 x more active than YL-03709B-A against the yeast *Candida albicans* and the filamentous fungus *Aspergillus fumigatus*, in view of the information for YL-

03709B-A reported in Japanese patent application 06-27802. Basiliskamide B also is significantly more active against these organisms than YL-03709B-A.

Antimycobacterial Activity of the Basiliskamides: The Basiliskamides were tested for antimycobacterial activity using a standardized agar dilution method (Underlied, CB and Salfinger, S. 1995. In Manual of Clinical Microbiology, Murray, Baron, Pfaller, Tenover, Yolken (Eds.), ASM Press, page 1395-1404). Activity of basiliskamide A, basiliskamide B, and acylated derivatives of the two compounds were tested for activity against *Mycobacterium tuberculosis*, the cause of tuberculosis, and *Mycobacterium avium-intracellulare*, an important cause of mycobacterial infections in immunocompromized patients such as those with AIDS. The results are shown in Table 6.

Table 6. Antimycobacterial activity of the Basiliskamides

15

10

5

## Minimal Inhibitory Concentration (µg/ml)

20	
Basiliskamide A 25 100 Basiliskamide B 50 > 100 Acylated A (2-16) > 100 > 100  25 Acylated B (2-16) > 100 ≥ 50	

These results indicate that basiliskamide A and B each have activity against *M. tuberculosis*. Basiliskamide A has activity against *M. avium-intracellulare*, but basiliskamide B appears to be relatively inactive against this organism. Decreasing the number of carbons in the backbone of the molecule may increase activity. Such is the case with the increased antifungal activity of the basiliskamides (A and B) that have one less carbon in the molecule's backbone than YL-03709B-A.

35

Cytotoxicity Testing of Basiliskamide A: Serial dilutions of basiliskamide A were prepared in cell culture medium and tested for toxicity for normal human fibroblast cells and for human tumor cell line. The effect of basiliskamide (basil) was compared to that of the known antifungal compound amphotericin B (ampho). The appearance of the cells was assessed after 48 hours exposure to the compounds. The results are shown in Table 7. Basiliskamide produced no cytotoxicity for normal human fibroblast cells at concentrations less than  $100 \mu g/ml$  compared to amphotericin B (ampho) which was toxic at concentrations as low as  $25 \mu g/ml$ . Against human tumor cells, basiliskamide showed minor toxicity at concentrations above  $3 \mu g/ml$ . These findings suggested that basiliskamide is less toxic for normal human cells than the widely used amphotericin B.

Table 7 Basiliskamide Cytoxicity Testing

1	-
	~

20

10

5

Cytopathic Effect (48 hours)					
Human Fibroblast Human Tumor					
Conc (µg/ml	Basil	Ampho	Basil	Ampho	
100	2	4	4	4	
50	0	2	1	3	
25	0	2	1	1	
12.5	0	1	1	0	
6.25	0	0	2	0	
3.12	0	0	2	0	
1.57	0	0	0	0	
0.78	0	0	0	0	
0	0	0	0	0	

25

- 1 slight change in morphology vs. control
- 2 occasional rounding, vacuolization, or granularity
- 3 rounding, vacuolization, detachment of 50% of cells
- 35 4 destruction of monolayer

All publications, patents and patent applications referred to herein are hereby incorporated by reference. While this invention has been described according to particular embodiments and by reference to certain examples, it will be apparent to those of skill in the art that variations and modifications of the invention as described herein fall within the spirit and scope of the attached claims.

#### 5 WE CLAIM:

1. A compound or a physiologically acceptable salt thereof, wherein the compound has the formula:

10

$$Z^{1}$$
 $R_{1}$ 
 $R_{2}$ 
 $R_{3}$ 
 $R_{4}$ 

15

wherein:

R<sub>1</sub> and R<sub>2</sub> are the same or different and are independently H or R;

R is a structural fragment having a saturated or unsaturated linear, branched, or cyclic, skeleton containing one to ten carbon atoms in which the carbon atoms may be optionally substituted with a substituent selected from the group consisting of: -OH; =O; -OR<sub>5</sub>; -O<sub>2</sub>CR<sub>5</sub>, -SH; -SR<sub>5</sub>; -SOCR<sub>5</sub>; -NH<sub>2</sub>; -NHR<sub>5</sub>; -NH(R<sub>5</sub>)<sub>2</sub>; -NHCOR<sub>5</sub>; NRCOR<sub>5</sub>; -I; -Br; -CI; -F; -CN; -CO<sub>2</sub>H; -CO<sub>2</sub>R<sub>5</sub>; -CHO; -COR<sub>5</sub>; -CONH<sub>2</sub>; -CONHR<sub>5</sub>; -CON(R<sub>5</sub>)<sub>2</sub>; -COSH; -COSR<sub>5</sub>; -NO<sub>2</sub>; -SO<sub>3</sub>H; -SOR<sub>5</sub>; and -SO<sub>2</sub>R<sub>5</sub>, wherein R<sub>5</sub> is a linear, branched or cyclic, one to ten carbon saturated or unsaturated alkyl group;

 $R_3$  and  $R_4$  are different and are independently selected from the groups consisting of OH,

20

25

(b) -O-Z-AI

wherein,

Z<sup>1</sup> and Z are linear or branched, saturated or unsaturated, one to ten carbon fragments
 optionally substituted with Y;

Ar is a monocyclic, bicyclic or tricyclic, fully or partially aromatic system containing five or six membered carbocyclic or, oxygen, nitrogen or sulphur containing heterocyclic rings, optionally substituted with R or Y;

Y is selected from the group consisting of: H; =O, -OH; -OR; -O<sub>2</sub>CR; -SH: -SR; -SOCR; -NH<sub>2</sub>; -NHR; -NH(R)<sub>2</sub>; -NHCOR; NRCOR; -I; -Br; -Cl; -F; -CN- -CO<sub>2</sub>H; -CO<sub>2</sub>R; -CHO; -COR; -CONH<sub>2</sub>; -CONHR; -CON(R)<sub>2</sub>; -COSH; -COSR; -NO<sub>2</sub>: -SO<sub>3</sub>H; -SOR; -SO<sub>2</sub>R; and, -O- (epoxide);

W is H or R;

with the provisos that when W is H,  $R_2$  is not H; when  $R_2$  is  $CH_3$ , W is not n-propyl; and, one of  $R_3$  and  $R_4$  is (a) or (b) and another of  $R_3$  and  $R_4$  is OH.

2. The compound or physiologically acceptable salt thereof of claim 1 having the stereoisomeric form:

30 Z

$$Z^{1}$$
 $R_{1}$ 
 $R_{2}$ 
 $W$ 
 $R_{3}$ 
 $R_{4}$ 

The compound or physiologically acceptable salt thereof of claim 1 or 2 wherein Z<sup>1</sup> is a linear or branched, saturated or unsaturated one to eight carbon carbonyl optionally substituted with a substituent selected from the group consisting of: NH<sub>2</sub>, NHR, NR<sub>2</sub>, OH, OR, SH, SR, H and CF<sub>3</sub>, wherein R is as defined.

10

4. A compound or a physiologically acceptable salt thereof, wherein the compound has the formula:

15

20

wherein:

a single, double or triple bond exists between one or more of: C-2 and C-3; C-3 and C-4; C-4 and C-5; and, C-5 and C-6;

25

X is NH<sub>2</sub>, NHR, NR<sub>2</sub>, OH, OR, SH, SR, H, or CF<sub>3</sub>;

R is a structural fragment having a saturated or unsaturated linear, branched, or cyclic. skeleton containing one to ten carbon atoms in which the carbon atoms may be optionally substituted with a substituent selected from the group consisting of: -OH: =O; -OR<sub>5</sub>; -O<sub>2</sub>CR<sub>5</sub>, -SH; -SR<sub>5</sub>; -SOCR<sub>5</sub>; -NH<sub>2</sub>; -NHR<sub>5</sub>; -NH(R<sub>5</sub>)<sub>2</sub>; -NHCOR<sub>5</sub>; NRCOR<sub>5</sub>; -I; -Br; -Cl; -F; -CN; -CO<sub>2</sub>H; -CO<sub>2</sub>R<sub>5</sub>; -CHO; -COR<sub>5</sub>; -CONH<sub>2</sub>; -CONHR<sub>5</sub>: -CON(R<sub>5</sub>)<sub>2</sub>; -COSH; -COSR<sub>5</sub>; -NO<sub>2</sub>; -SO<sub>3</sub>H; -SOR<sub>5</sub>: and -SO<sub>2</sub>R<sub>5</sub>, wherein R<sub>5</sub> is a linear, branched or cyclic, one to ten carbon saturated or unsaturated alkyl group;

35

30

R<sub>1</sub> and R<sub>2</sub> are the same or different and are independently H or R;

20

R<sub>3</sub> and R<sub>4</sub> are different and are selected from the group consisting of: OH,

and

wherein, Z is a linear or branched, saturated or unsaturated, one to ten carbon fragment optionally substituted with Y;

Ar is a monocyclic, bicyclic or tricyclic, fully or partially aromatic system containing five or six membered carbocyclic or, oxygen, nitrogen or sulphur containing heterocyclic rings, optionally substituted with R or Y:

Y is selected from the group consisting of: H; =O. -OH; -OR; -O<sub>2</sub>CR; -SH; -SR; -SOCR; -NH<sub>2</sub>; -NHR; -NH(R)<sub>2</sub>; -NHCOR; NRCOR: -I; -Br; -Cl; -F; -CN- -CO<sub>2</sub>H; -CO<sub>2</sub>R; -CHO; -COR; -CONH<sub>2</sub>; -CONHR; -CON(R)<sub>2</sub>; -COSH; -COSR; -NO<sub>2</sub>; -SO<sub>3</sub>H; -SOR; -SO<sub>2</sub>R; and, -O- (epoxide);

with the proviso that one of R<sub>3</sub> and R<sub>4</sub> is (a) or (b), and another of R<sub>3</sub> and R<sub>4</sub> is OH.

5. The compound or physiologically acceptable salt thereof of claim 4 having the structure:

$$X \xrightarrow{Q} X \xrightarrow{R_1} \xrightarrow{R_2} X$$

6. The compound or physiologically acceptable salt thereof of claim 4, having the structural and stereoisomeric form:

10

$$X$$
 $R_1$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 

15

- 7. The compound or physiological salt thereof of any one of claims 4-6, wherein R<sub>1</sub> and R<sub>2</sub> are independently H or CH<sub>3</sub>.
  - 8. The compound or physiological salt thereof of any one of claims 4-7, wherein R<sub>3</sub> is (a).

- 9. The compound or physiological salt thereof of any one of claims 4-8, wherein X is NH<sub>2</sub>.
- The compound or physiological salt thereof of any one of claims 4-9, wherein R<sub>3</sub> at C<sub>7</sub> is (a) and R<sub>3</sub> at C<sub>9</sub> is OH.
  - 11. The compound or physiological salt thereof of any one of claims 4-9, wherein R<sub>3</sub> at C<sub>7</sub> is OH and R<sub>3</sub> at C<sub>9</sub> is (a).

- 5 12. A compound according to claim 4, wherein the compound is Basiliskamide A substantially free of cellular contaminants.
  - 13. A compound according to claim 4, wherein the compound is Basiliskamide B substantially free of cellular contaminants.
  - 14. A pharmaceutical composition comprising a compound or physiological salt thereof of any one of claims 1-13, and a pharmaceutically acceptable carrier.
- 15. The use of a compound or physiological salt thereof of any one of claims 1-13.

  as an antifungal agent.
  - 16. The use of a compound or physiological salt thereof of any one of claims 1-3. as an antimycobacterial agent.

Antibiotic polyketide compounds are provided having the formula

10

$$Z^{1}$$
 $R_{1}$ 
 $R_{2}$ 
 $R_{3}$ 
 $R_{4}$ 

15

wherein:

R<sub>1</sub> and R<sub>2</sub> are the same or different and are independently H or R;

R is a structural fragment having a saturated or unsaturated linear, branched, or cyclic. skeleton containing one to ten carbon atoms in which the carbon atoms may be optionally substituted with a substituent selected from the group consisting of: -OH; =O; -OR<sub>5</sub>; -O<sub>2</sub>CR<sub>5</sub>, -SH; -SR<sub>5</sub>; -SOCR<sub>5</sub>; -NH<sub>2</sub>; -NHR<sub>5</sub>; -NH(R<sub>5</sub>)<sub>2</sub>; -NHCOR<sub>5</sub>; NRCOR<sub>5</sub>; -I; -Br; -Cl; -F; -CN; -CO<sub>2</sub>H; -CO<sub>2</sub>R<sub>5</sub>; -CHO; -COR<sub>5</sub>; -CONH<sub>2</sub>; -CONHR<sub>5</sub>; -CON(R<sub>5</sub>)<sub>2</sub>; -COSH; -COSR<sub>5</sub>; -NO<sub>2</sub>; -SO<sub>3</sub>H; -SOR<sub>5</sub>; and -SO<sub>2</sub>R<sub>5</sub>, wherein R<sub>5</sub> is a linear, branched or cyclic, one to ten carbon saturated or unsaturated alkyl group;

R<sub>3</sub> and R<sub>4</sub> are different and are independently selected from the groups consisting of OH,

(b) 
$$-O-Z-Ar$$

wherein.

Z<sup>1</sup> and Z are linear or branched, saturated or unsaturated, one to ten carbon fragments
 optionally substituted with Y;

Ar is a monocyclic, bicyclic or tricyclic, fully or partially aromatic system containing five or six membered carbocyclic or, oxygen, nitrogen or sulphur containing heterocyclic rings, optionally substituted with R or Y;

15

Y is selected from the group consisting of: H; =O, -OH; -OR; -O<sub>2</sub>CR; -SH; -SR; -SOCR; -NH<sub>2</sub>; -NHR; -NH(R)<sub>2</sub>; -NHCOR; NRCOR; -I; -Br; -CI; -F; -CN- -CO<sub>2</sub>H; -CO<sub>2</sub>R; -CHO; -COR; -CONH<sub>2</sub>; -CONHR; -CON(R)<sub>2</sub>; -COSH; -COSR; -NO<sub>2</sub>; -SO<sub>3</sub>H: -SOR; -SO<sub>2</sub>R; and, -O- (epoxide);

20

W is H or R;

with the provisos that when W is H,  $R_2$  is not H; when  $R_2$  is  $CH_3$ , W is not n-propyl: and, one of  $R_3$  and  $R_4$  is (a) or (b) and another of  $R_3$  and  $R_4$  is OH.